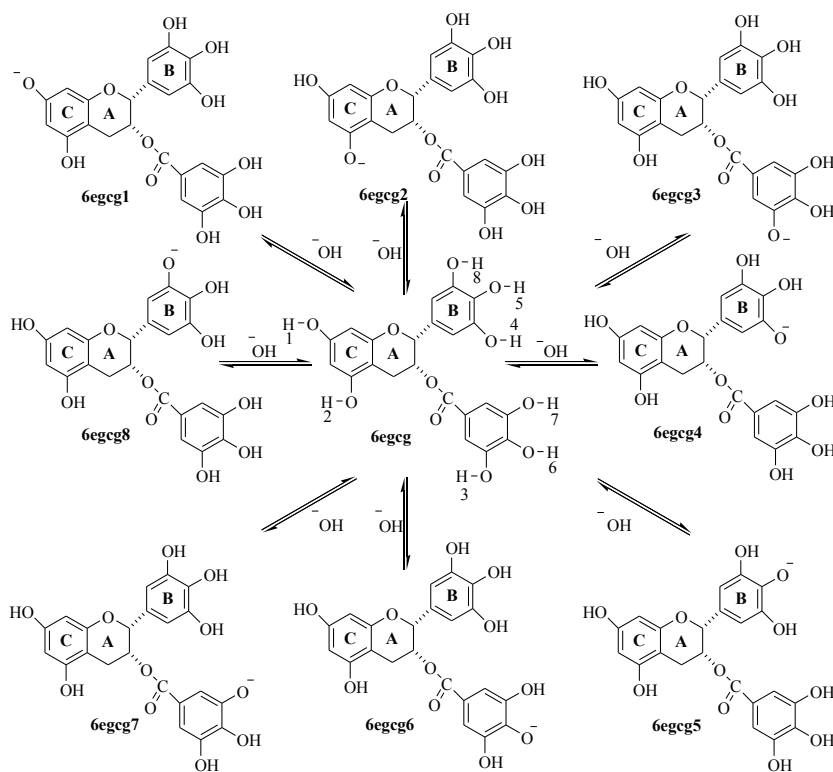


Epigallo catechin  
(5egc)



Epigallo catechin gallate  
(6egcg)

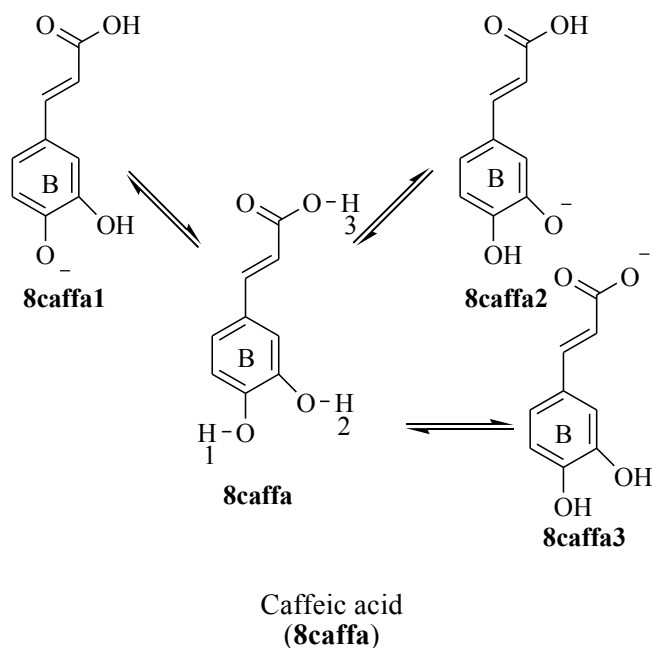
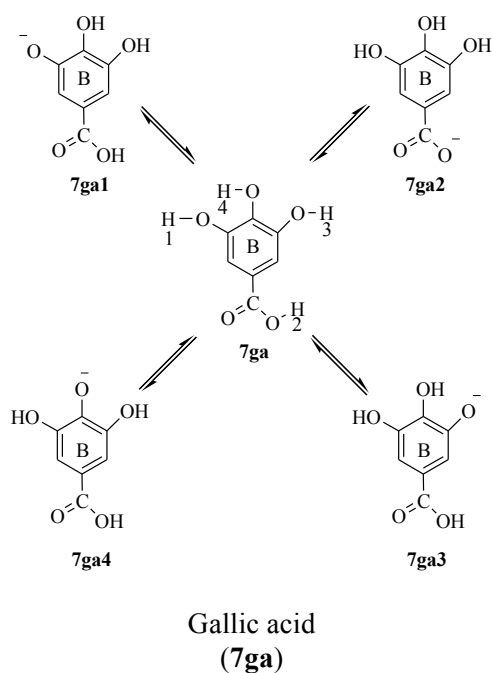


Figure 1. The schematic illustrations for the protonation processes of the tea catechins and polyphenols

The self-computed thermodynamic data was used in predicting the  $pK_a$  values of various species, using Eq. 5 where  $\delta\Delta G_{(BH)}$  is the standard free energy charge for the reaction (1).

$$pK_{a(BH)} = \delta\Delta G_{(BH)} / (2.303RT) \quad (5)$$

$(\Delta G_f)$  and  $(\Delta G)$ , standard free energy symbols, represent thermokinetic and thermodynamic data, respectively.

In the theoretical calculations, both aqueous and blood phases were used for determining the activities of catechins and polyphenols in the human metabolism to calculate the most reactive component among these molecules in this study (Table 1).

Table 1. Certain physicochemical parameters of the studied molecules and mono anion forms in aqueous (T=298 K,  $\epsilon=78.4$ ) and blood (T=310 K,  $\epsilon=58$ ) phases calculated by the PM6 method

| Molecules <sup>a</sup> | $\Delta H_f^b$                             | $\Delta H^c$ | $\Delta S^d$ | $\Delta G_f^e$ | $\Delta G^f$ | $\Delta H_f^b$                          | $\Delta H^c$ | $\Delta S^d$ | $\Delta G_f^e$ | $\Delta G^f$ |
|------------------------|--|--------------|--------------|----------------|--------------|---|--------------|--------------|----------------|--------------|
|                        | Neutral (BH)-aqueous phase                 |              |              |                |              | Neutral (BH)-blood phase                |              |              |                |              |
| 1c                     | -239.801                                   | 13074.095    | 145.929      | -283.288       | -30.413      | -238.680                                | 14026.146    | 148.743      | -283.005       | -30.299      |
| 2ec                    | -243.375                                   | 12963.362    | 143.525      | -286.145       | -29.807      | -242.253                                | 13920.070    | 146.709      | -285.972       | -29.799      |
| 3cg                    | -385.150                                   | 20049.185    | 204.660      | -446.139       | -40.939      | -383.467                                | 21486.057    | 209.482      | -445.892       | -40.939      |
| 4ecg                   | -384.901                                   | 20054.308    | 202.853      | -445.351       | -40.396      | -383.225                                | 21493.565    | 207.065      | -444.930       | -40.212      |
| 5egc                   | -287.474                                   | 14022.126    | 154.391      | -333.482       | -31.986      | -286.279                                | 15030.073    | 157.769      | -333.294       | -31.985      |
| 6egcg                  | -429.134                                   | 20933.933    | 213.825      | -492.854       | -42.786      | -427.381                                | 22427.901    | 219.229      | -492.711       | -42.902      |
| 7ga                    | -206.948                                   | 8314.803     | 109.713      | -239.643       | -24.380      | -206.271                                | 8878.540     | 111.563      | -239.517       | -24.367      |
| 8caffa                 | -151.129                                   | 8763.721     | 113.538      | -184.963       | -25.071      | -150.377                                | 9367.799     | 115.560      | -184.814       | -25.069      |
|                        | Mono anion (B <sup>-</sup> )-aqueous phase |              |              |                |              | Mono anion(B <sup>-</sup> )-blood phase |              |              |                |              |
| 1c1                    | -335.892                                   | 12978.431    | 146.222      | -379.466       | -30.596      | -334.392                                | 13920.978    | 149.352      | -378.899       | -30.586      |
| 1c2                    | -336.078                                   | 12990.150    | 146.017      | -379.591       | -30.523      | -334.604                                | 13922.108    | 148.973      | -378.998       | -30.472      |
| 1c3                    | -316.509                                   | 12970.805    | 145.877      | -359.980       | -30.501      | -315.004                                | 13906.298    | 148.995      | -359.404       | -30.494      |
| 1c4                    | -332.700                                   | 12844.879    | 143.511      | -375.466       | -29.921      | -331.201                                | 13775.755    | 146.629      | -374.896       | -29.920      |
| 1c5                    | -335.663                                   | 12989.469    | 146.859      | -379.427       | -30.775      | -334.226                                | 13927.600    | 150.158      | -378.973       | -30.819      |
| 2ec1                   | -335.881                                   | 12797.015    | 143.909      | -378.766       | -30.088      | -334.383                                | 13730.124    | 147.097      | -378.218       | -30.105      |
| 2ec2                   | -335.944                                   | 12801.506    | 143.548      | -378.721       | -29.976      | -334.462                                | 13742.350    | 146.606      | -378.151       | -29.946      |
| 2ec3                   | -316.557                                   | 12943.610    | 145.952      | -360.051       | -30.550      | -315.055                                | 13880.583    | 148.941      | -359.439       | -30.504      |
| 2ec4                   | -332.747                                   | 12954.548    | 146.501      | -376.404       | -30.703      | -331.250                                | 13890.080    | 148.437      | -375.484       | -30.344      |
| 2ec5                   | -335.812                                   | 12823.458    | 145.693      | -379.228       | -30.593      | -334.387                                | 13757.568    | 148.899      | -378.759       | -30.614      |
| 3cg1                   | -471.355                                   | 19479.829    | 197.371      | -530.172       | -39.337      | -469.421                                | 20893.799    | 201.481      | -529.462       | -39.148      |
| 3cg2                   | -476.540                                   | 19468.091    | 198.848      | -535.797       | -39.788      | -474.672                                | 20874.055    | 202.513      | -535.021       | -39.475      |
| 3cg3                   | -483.136                                   | 18992.383    | 187.953      | -539.146       | -37.017      | -481.198                                | 20408.536    | 192.634      | -538.603       | -36.997      |
| 3cg4                   | -472.481                                   | 19509.323    | 195.468      | -530.731       | -38.740      | -470.575                                | 20918.620    | 200.242      | -530.247       | -38.754      |
| 3cg5                   | -478.426                                   | 19195.166    | 192.215      | -535.706       | -38.085      | -476.563                                | 20597.878    | 196.732      | -535.189       | -38.028      |
| 3cg6                   | -475.456                                   | 19353.347    | 193.156      | -533.016       | -38.207      | -473.619                                | 20760.485    | 198.116      | -532.658       | -38.278      |
| 3cg7                   | -486.034                                   | 19257.003    | 192.847      | -543.503       | -38.212      | -484.188                                | 20667.400    | 197.685      | -543.098       | -38.243      |
| 4ecg1                  | -471.687                                   | 19498.418    | 196.943      | -530.376       | -39.190      | -469.730                                | 20914.453    | 201.584      | -529.802       | -39.157      |
| 4ecg2                  | -477.680                                   | 19393.601    | 194.409      | -535.614       | -38.540      | -475.789                                | 20806.886    | 199.188      | -535.147       | -38.551      |
| 4ecg3                  | -479.132                                   | 19651.262    | 197.093      | -537.866       | -39.082      | -477.218                                | 21062.056    | 201.719      | -537.330       | -39.050      |
| 4ecg4                  | -476.620                                   | 19402.424    | 197.228      | -535.394       | -39.371      | -474.713                                | 20804.890    | 201.806      | -534.851       | -39.333      |
| 4ecg5                  | -474.961                                   | 19360.941    | 192.829      | -532.424       | -38.102      | -473.112                                | 20766.840    | 197.526      | -531.975       | -38.096      |
| 4ecg6                  | -479.268                                   | 19455.850    | 198.146      | -538.315       | -39.592      | -477.386                                | 20862.539    | 202.368      | -537.692       | -39.443      |
| 4ecg7                  | -478.183                                   | 18949.967    | 186.040      | -533.623       | -36.490      | -476.252                                | 20361.693    | 190.747      | -533.095       | -36.481      |
| 5egc1                  | -380.671                                   | 13984.596    | 152.897      | -426.234       | -31.579      | -379.114                                | 14986.702    | 156.187      | -425.658       | -31.557      |
| 5egc2                  | -380.671                                   | 13984.596    | 152.897      | -426.234       | -31.579      | -379.068                                | 15032.116    | 155.367      | -425.367       | -31.267      |
| 5egc3                  | -361.644                                   | 13460.179    | 147.480      | -405.593       | -30.489      | -360.090                                | 14456.724    | 150.877      | -405.051       | -30.505      |
| 5egc4                  | -382.393                                   | 13775.795    | 151.444      | -427.523       | -31.354      | -380.899                                | 14772.058    | 154.927      | -427.067       | -31.396      |
| 5egc5                  | -377.537                                   | 13902.378    | 152.519      | -422.988       | -31.548      | -375.998                                | 14899.220    | 155.877      | -422.449       | -31.552      |
| 5egc6                  | -380.031                                   | 13778.410    | 151.625      | -425.215       | -31.406      | -378.471                                | 14771.054    | 155.035      | -424.671       | -31.429      |

Table 1. (Continued) Certain physicochemical parameters of the studied molecules and mono anion forms in aqueous (T=298 K,  $\epsilon=78.4$ ) and blood (T=310 K,  $\epsilon=58$ ) phases calculated by the PM6 method

|                                   |          |           |         |          |         |          |           |         |          |         |
|-----------------------------------|----------|-----------|---------|----------|---------|----------|-----------|---------|----------|---------|
| <b>6egcg1</b>                     | -520.014 | 20707.281 | 207.079 | -581.723 | -41.002 | -517.950 | 22188.203 | 212.635 | -581.315 | -41.177 |
| <b>6egcg2</b>                     | -521.136 | 20258.753 | 200.287 | -580.822 | -39.427 | -519.197 | 21717.718 | 205.019 | -580.293 | -39.378 |
| <b>6egcg3</b>                     | -519.750 | 20021.323 | 198.268 | -578.834 | -39.062 | -517.736 | 21484.926 | 203.037 | -578.241 | -39.020 |
| <b>6egcg4</b>                     | -514.503 | 20557.285 | 202.014 | -574.703 | -39.643 | -512.463 | 22033.251 | 207.186 | -574.204 | -39.708 |
| <b>6egcg5</b>                     | -523.320 | 20288.472 | 200.768 | -583.149 | -39.540 | -521.366 | 21754.615 | 205.718 | -582.670 | -39.549 |
| <b>6egcg6</b>                     | -524.640 | 20322.005 | 201.958 | -584.824 | -39.862 | -522.641 | 21792.862 | 207.019 | -584.333 | -39.899 |
| <b>6egcg7</b>                     | -525.130 | 20379.775 | 204.348 | -586.026 | -40.516 | -523.172 | 21842.428 | 208.134 | -585.196 | -40.181 |
| <b>6egcg8</b>                     | -531.073 | 20353.635 | 202.515 | -591.422 | -39.996 | -529.159 | 21832.769 | 208.937 | -591.422 | -40.430 |
| <b>7ga1</b>                       | -304.427 | 7873.122  | 105.755 | -335.942 | -23.642 | -303.408 | 8405.959  | 107.505 | -335.444 | -23.630 |
| <b>7ga2</b>                       | -306.558 | 7745.196  | 104.684 | -337.754 | -23.451 | -305.472 | 8282.691  | 106.512 | -337.213 | -23.458 |
| <b>7ga3</b>                       | -303.076 | 7871.621  | 107.744 | -335.184 | -24.236 | -302.016 | 8407.490  | 109.444 | -334.630 | -24.207 |
| <b>7ga4</b>                       | -304.004 | 7834.405  | 106.641 | -335.783 | -23.944 | -302.957 | 8367.778  | 108.325 | -335.238 | -23.913 |
| <b>8caffa1</b>                    | -246.797 | 8286.870  | 111.706 | -280.085 | -25.001 | -245.783 | 8855.535  | 113.573 | -279.628 | -24.989 |
| <b>8caffa2</b>                    | -241.906 | 8134.321  | 109.337 | -274.488 | -24.448 | -240.851 | 8700.298  | 111.211 | -273.992 | -24.441 |
| <b>8caffa3</b>                    | -250.267 | 8394.615  | 111.825 | -283.591 | -24.929 | -249.163 | 8969.410  | 113.708 | -283.048 | -24.915 |
| <b>H<sub>2</sub>O</b>             | -61.357  | 2373.258  | 45.108  | -74.799  | -11.069 | -61.136  | 2376.435  | 45.005  | -74.547  | -11.035 |
| <b>H<sub>3</sub>O<sup>+</sup></b> | 48.753   | 2383.074  | 46.419  | 34.920   | -11.450 | 58.582   | 2427.178  | 44.734  | 45.251   | -10.904 |

<sup>a</sup>: The last number of the symbols of the molecules represents leaving -H atom numbers marked in Fig. 1

<sup>b</sup>  $\Delta H_f$ : Heat of formation;

<sup>c</sup>  $\Delta H$ : Enthalpy; and

<sup>d</sup>  $\Delta S$ : Entropy were calculated by MOPAC2009.

<sup>e</sup>  $\Delta G_f$ : Formation Gibbs free energy was calculated by the equation of  $\Delta G_f = \Delta H_f - T\Delta S$

<sup>f</sup>  $\Delta G$ : Gibbs free energy was calculated by  $\Delta G = \Delta H - T\Delta S$

## 2.1 Gibbs Free Energy ( $\Delta G$ and $\Delta G_f$ )

As can be seen from Table 1, the order of stabilities of these molecules depends on their Gibbs Free Energies ( $\Delta G$  and  $\Delta G_f$ ) in the blood phase being the same as those in the aqueous phase. It is observed that any molecule having more of the -OH group has more stability and is less reactive. It is thought that the reason for this originates from a possible solvation. When the stabilities in the aqueous and blood phases are examined it is realized that they are close to each other in two phases but the stabilities of all the molecules are higher in the aqueous phase than those in the blood phase. This case results from the higher solvation strength of the water. The best reactive structures are 8caffa and 7ga among these molecules. In addition to these findings, the  $\Delta G$  order for 8caffa and the 7ga are different from those for the  $\Delta G_f$  order. It is thought that 8caffa in the order of  $\Delta G$  is dominant kinetically and 7ga in the order of  $\Delta G_f$  is dominant thermodynamically.

Certain catechin molecules have epimers. In fact, this situation which has its origins in molecular geometry is crucial for human metabolism. In the aqueous and blood phases, 2ec (including 5 -OH group) and 3cg (including 7 -OH group) are more stable than 1c (having 5 -OH) and 4ecg (having 7 -OH group), respectively.

In Fig. 1 an 'A' ring represents a saturated six carbon ring for the studied molecules and this ring plays an important role from the point of view of stability in the configuration. The 'A' ring does not lie in one plane. Thus, the ring prefers the most stable conformation. 2ec and 4ecg, epimer couples, placed on the more stable position than 1c and 3cg. 1c and 3cg placed in the steric affected position. Even if the energies of these four molecules are not too different from each other, 3cg and 4ecg are more stable than 1c and 2ec and their reactivities are less than 1c and 2ec.



**1c** (having 5 –OH group), **2ec** (having 5-OH group), **5egc** (having 6 –OH group), **3cg** (having 7 –OH group), **4ecg** (having 7 –OH group) and **6egcg** (having 8 –OH group) have the same structure. But, the ‘**B**’ ring in **5egc** and **6egcg** has one more –OH than the others. The stability order is **6egcg**>**3cg**>**4ecg**>**5egc**>**2ec**>**1c** in both the aqueous and the blood phases. **6egcg**, **3cg** and **4ecg** are more stable, less reactive and have less biologically activity than **5egc**, **2ec** and **1c**.

**7ga** and **8caffa** are the most reactive molecules among the studied compounds. The structures of these molecules are totally different from the other molecules. **7ga** and **8caffa** include the least –OH group among the studied catechins. Since they have one ring, their reactivities are higher than the catechins.

In conclusion, it is thought that not only the number of –OH group but also the geometry of the tea components has an important role in the reactivities of these compounds in human metabolism.

Table 2. Certain physical parameters of the studied molecules in the aqueous (T=298 K, ε=78.4) and blood (T=310 K, ε=58) phases

| Molecules     | HOMO          | LUMO   | N <sup>a</sup> | D <sup>b</sup> | HOMO        | LUMO   | N <sup>a</sup> | D <sup>b</sup> |
|---------------|---------------|--------|----------------|----------------|-------------|--------|----------------|----------------|
|               | Aqueous phase |        |                |                | Blood phase |        |                |                |
| <b>1c</b>     | -9.066        | -0.322 | -8.744         | 5.764          | -9.064      | -0.320 | -8.744         | 5.747          |
| <b>2ec</b>    | -9.125        | -0.434 | -8.691         | 4.587          | -9.125      | -0.434 | -8.691         | 4.573          |
| <b>3cg</b>    | -9.175        | -1.287 | -7.888         | 7.193          | -9.175      | -1.286 | -7.889         | 7.172          |
| <b>4ecg</b>   | -9.163        | -1.312 | -7.851         | 6.705          | -9.161      | -1.312 | -7.849         | 6.689          |
| <b>5egc</b>   | -9.449        | -0.595 | -8.854         | 5.127          | -9.448      | -0.595 | -8.853         | 5.115          |
| <b>6egcg</b>  | -9.457        | -1.290 | -8.167         | 4.963          | -9.454      | -1.288 | -8.166         | 4.951          |
| <b>7ga</b>    | -9.444        | -1.272 | -8.172         | 5.707          | -9.443      | -1.271 | -8.172         | 5.690          |
| <b>8caffa</b> | -9.090        | -1.265 | -7.825         | 6.664          | -9.091      | -1.265 | -7.826         | 6.652          |

<sup>a</sup>N=HOMO-LUMO

<sup>b</sup>D: Dipole moment

## 2.2 Dipole Moment

When the dipole moment (**D**) of the molecules was investigated, no important correlation could be observed between the molecule structures and reactivities for all of the studied molecules (Table 2). It is thought that the dipole moment does not give any important information from the viewpoint of the reactivity for these kinds of molecules in the present study.

## 2.3 Nucleophilicity

When the nucleophilicities (**N** =HOMO-LUMO) of the molecules are investigated, the N value order is given in the following order: **5egc** (including the 6-OH group)>**1c** (including 5 –OH group) >**2ec** (including 5 –OH group) and **6egcg** (including 8 -OH group)>**3cg** (including 7 -OH group) >**4ecg** (including 7–OH group). But, the N values of **7ga** and **8caffa** are close to catechins. It is therefore thought that the nucleophilicity does not show an important effect on determining the reactivity.

## 2.4 pK<sub>a</sub> (PM6, Thermodynamic, Thermokinetic, Marvinbeans, Sparc)

The pK<sub>a</sub> values of the tea compounds were computed by PM6 (automatically computed by MOPAC2009), Thermodynamic (ΔG), Thermokinetic (ΔG<sub>f</sub>), Marvinbeans and Sparc methods (Table 3-5).

As it can be seen from Fig. 2 and Fig. 3, the pK<sub>a</sub> values, calculated using different methods [PM6, ΔG<sub>f</sub> (Thermokinetic), ΔG (Thermodynamic), Marvinbeans and Sparc], were compared with those of

the experimental ones in both the aqueous and the blood phases. The  $pK_a$  values were compared by computation with different methods in both aqueous and blood phases; the best correlations could be observed between PM6 and experimental  $pK_a$  values for **6egcg** and **8caffa**; PM6- $\Delta G_f$  for **4ecg**; Marvin-experimental for **1c** and **2ec**;  $\Delta G_f$ -Sparc for **3cg**,  $\Delta G_f$ - $\Delta G$  for **5egc** and Marvinbeans-Sparc for **7ga** in the aqueous phase; and Marvinbeans-experimental for **1c** and **2ec**;  $\Delta G_f$ -Marvinbeans for **3cg**;  $\Delta G$ -experimental for **4ecg**;  $\Delta G_f$ -Sparc for **5egc**;  $\Delta G$ -Sparc for **6egcg**; and Marvinbeans-Sparc for **7ga** and  $\Delta G_f$ -experimental for **8caffa** in the blood phase.

A good correlation could not be observed among the  $pK_a$  values for the similar acidic proton belonging to the -OH in the molecules for all calculation methods [PM6,  $\Delta G_f$  (Thermokinetic),  $\Delta G$  (Thermodynamic), Marvinbeans and Sparc] (Table 5).

Positive meaningful correlations ( $R^2 > 0.9999$ ) were observed between thermokinetic  $pK_a$  values both in blood and aqueous phases for **1c**, **2ec**, **3cg**, **4ecg**; **5egc**, **6egcg** and **7ga** (any table or figure was not given here).

Table 3. Thermokinetic  $pK_a$  values of the molecules in the aqueous (T=298 K,  $\epsilon=78.4$ ) and blood (T=310 K,  $\epsilon=58$ ) phases

| Protonation process <sup>a</sup> | $\Delta G_f(B)^b$ | $\Delta G_f(BH)^c$ | $\delta\Delta G_f(BH)^d$ | $pK_{af}(BH)^e$ | $\Delta G_f(B)^b$ | $\Delta G_f(BH)^c$ | $\delta\Delta G_f(BH)^d$ | $pK_{af}(BH)^e$ |
|----------------------------------|-------------------|--------------------|--------------------------|-----------------|-------------------|--------------------|--------------------------|-----------------|
|                                  | Aqueous phase     |                    |                          |                 | Blood phase       |                    |                          |                 |
| <b>1c-1c1</b>                    | -379.466          | -283.288           | 13.541                   | 9.930           | -378.899          | -283.005           | 23.904                   | 17.530          |
| <b>1c-1c2</b>                    | -379.591          | -283.288           | 13.416                   | 9.838           | -378.998          | -283.005           | 23.805                   | 17.457          |
| <b>1c-1c3</b>                    | -359.980          | -283.288           | 33.027                   | 24.219          | -359.404          | -283.005           | 43.399                   | 31.825          |
| <b>1c-1c4</b>                    | -375.466          | -283.288           | 17.541                   | 12.863          | -374.896          | -283.005           | 27.907                   | 20.465          |
| <b>1c-1c5</b>                    | -379.427          | -283.288           | 13.580                   | 9.958           | -378.973          | -283.005           | 23.830                   | 17.475          |
| <b>2ec-2ec1</b>                  | -378.766          | -286.145           | 17.098                   | 12.539          | -378.218          | -285.972           | 27.552                   | 20.205          |
| <b>2ec-2ec2</b>                  | -378.721          | -286.145           | 17.143                   | 12.571          | -378.151          | -285.972           | 27.620                   | 20.254          |
| <b>2ec-2ec3</b>                  | -360.051          | -286.145           | 35.814                   | 26.263          | -359.439          | -285.972           | 46.331                   | 33.975          |
| <b>2ec-2ec4</b>                  | -376.404          | -286.145           | 19.460                   | 14.271          | -375.484          | -285.972           | 30.286                   | 22.209          |
| <b>2ec-2ec5</b>                  | -379.228          | -286.145           | 16.636                   | 12.199          | -378.759          | -285.972           | 27.011                   | 19.808          |
| <b>3cg-3cg1</b>                  | -530.172          | -446.139           | 25.686                   | 18.836          | -529.462          | -445.892           | 36.228                   | 26.567          |
| <b>3cg-3cg2</b>                  | -535.797          | -446.139           | 20.061                   | 14.711          | -535.021          | -445.892           | 30.670                   | 22.491          |
| <b>3cg-3cg3</b>                  | -539.146          | -446.139           | 16.712                   | 12.255          | -538.603          | -445.892           | 27.087                   | 19.864          |
| <b>3cg-3cg4</b>                  | -530.731          | -446.139           | 25.127                   | 18.426          | -530.247          | -445.892           | 35.443                   | 25.991          |
| <b>3cg-3cg5</b>                  | -535.706          | -446.139           | 20.151                   | 14.777          | -535.189          | -445.892           | 30.501                   | 22.367          |
| <b>3cg-3cg6</b>                  | -533.016          | -446.139           | 22.841                   | 16.750          | -532.658          | -445.892           | 33.033                   | 24.224          |
| <b>3cg-3cg7</b>                  | -543.503          | -446.139           | 12.355                   | 9.060           | -543.098          | -445.892           | 22.592                   | 16.567          |
| <b>4ecg-4ecg1</b>                | -530.376          | -445.351           | 24.694                   | 18.109          | -529.802          | -444.930           | 34.926                   | 25.612          |
| <b>4ecg-4ecg2</b>                | -535.614          | -445.351           | 19.456                   | 14.268          | -535.147          | -444.930           | 29.581                   | 21.693          |
| <b>4ecg-4ecg3</b>                | -537.866          | -445.351           | 17.205                   | 12.616          | -537.330          | -444.930           | 27.398                   | 20.091          |
| <b>4ecg-4ecg4</b>                | -535.394          | -445.351           | 19.676                   | 14.429          | -534.851          | -444.930           | 29.877                   | 21.909          |
| <b>4ecg-4ecg5</b>                | -532.424          | -445.351           | 22.646                   | 16.607          | -531.975          | -444.930           | 32.753                   | 24.019          |
| <b>4ecg-4ecg6</b>                | -538.315          | -445.351           | 16.755                   | 12.287          | -537.692          | -444.930           | 27.037                   | 19.826          |
| <b>4ecg-4ecg7</b>                | -533.623          | -445.351           | 21.447                   | 15.728          | -533.095          | -444.930           | 31.634                   | 23.198          |
| <b>5egc-5egc1</b>                | -426.234          | -333.482           | 16.967                   | 12.442          | -425.658          | -333.294           | 27.435                   | 20.118          |
| <b>5egc-5egc2</b>                | -426.234          | -333.482           | 16.967                   | 12.442          | -425.367          | -333.294           | 27.725                   | 20.331          |
| <b>5egc-5egc3</b>                | -405.593          | -333.482           | 37.609                   | 27.579          | -405.051          | -333.294           | 48.041                   | 35.229          |

Table 3. (Continued) Thermokinetic  $pK_a$  values of the molecules in the aqueous ( $T=298\text{ K}$ ,  $\epsilon=78.4$ ) and blood ( $T=310\text{ K}$ ,  $\epsilon=58$ ) phases

|                |          |          |        |        |          |          |        |        |
|----------------|----------|----------|--------|--------|----------|----------|--------|--------|
| 5egc-5egc4     | -427.523 | -333.482 | 15.678 | 11.497 | -427.067 | -333.294 | 26.025 | 19.085 |
| 5egc-5egc5     | -422.988 | -333.482 | 20.214 | 14.823 | -422.449 | -333.294 | 30.643 | 22.471 |
| 5egc-5egc6     | -425.215 | -333.482 | 17.986 | 13.190 | -424.671 | -333.294 | 28.421 | 20.842 |
| 6egcg-6egcg1   | -581.723 | -492.854 | 20.849 | 15.289 | -581.315 | -492.711 | 31.194 | 22.875 |
| 6egcg-6egcg2   | -580.822 | -492.854 | 21.751 | 15.951 | -580.293 | -492.711 | 32.217 | 23.625 |
| 6egcg-6egcg3   | -578.834 | -492.854 | 23.739 | 17.408 | -578.241 | -492.711 | 34.268 | 25.129 |
| 6egcg-6egcg4   | -574.703 | -492.854 | 27.870 | 20.437 | -574.204 | -492.711 | 38.305 | 28.089 |
| 6egcg-6egcg5   | -583.149 | -492.854 | 19.424 | 14.244 | -582.670 | -492.711 | 29.839 | 21.881 |
| 6egcg-6egcg6   | -584.824 | -492.854 | 17.749 | 13.016 | -584.333 | -492.711 | 28.176 | 20.662 |
| 6egcg-6egcg7   | -586.026 | -492.854 | 16.547 | 12.134 | -585.196 | -492.711 | 27.313 | 20.029 |
| 6egcg-6egcg8   | -591.422 | -492.854 | 11.150 | 8.177  | -591.422 | -492.711 | 21.087 | 15.463 |
| 7ga-7ga1       | -335.942 | -239.643 | 13.420 | 9.841  | -335.444 | -239.517 | 23.870 | 17.505 |
| 7ga-7ga2       | -337.754 | -239.643 | 11.608 | 8.512  | -337.213 | -239.517 | 22.102 | 16.208 |
| 7ga-7ga3       | -335.184 | -239.643 | 14.178 | 10.397 | -334.630 | -239.517 | 24.685 | 18.102 |
| 7ga-7ga4       | -335.783 | -239.643 | 13.579 | 9.957  | -335.238 | -239.517 | 24.077 | 17.656 |
| 8caffa-8caffa1 | -280.085 | -184.963 | 14.597 | 10.704 | -279.628 | -184.814 | 24.984 | 18.321 |
| 8caffa-8caffa2 | -274.488 | -184.963 | 20.194 | 14.809 | -273.992 | -184.814 | 30.620 | 22.454 |
| 8caffa-8caffa2 | -283.591 | -184.963 | 11.092 | 8.134  | -283.048 | -184.814 | 21.564 | 15.813 |

<sup>a</sup>: These symbols represent the protonation process for the molecules given in Fig.1

<sup>b</sup>  $\Delta G_f(B^-)$ : Formation Gibbs free energy of  $B^-$

<sup>c</sup>  $\Delta G_f(BH)$ : Formation Gibbs free energy of protonated  $BH$

<sup>d</sup>  $\delta\Delta G_f(BH)$ : Gibbs free energy calculated according to the Eq. 2 for the reaction (1)

<sup>e</sup>  $pK_{af}(BH)$ : Thermokinetic  $pK_a$  values calculated according to the Eq. 5

Table 4. Thermodynamic  $pK_a$  values of the molecules in the aqueous ( $T=298\text{ K}$ ,  $\epsilon=78.4$ ) and blood ( $T=310\text{ K}$ ,  $\epsilon=58$ ) phases

| Protonation process <sup>a</sup> | $\Delta G(B^-)^b$ | $\Delta G(BH)^c$ | $\delta\Delta G(BH)^d$ | $pK_{af}(BH)^e$ | Aqueous phase     |                  | Blood phase            |                 |
|----------------------------------|-------------------|------------------|------------------------|-----------------|-------------------|------------------|------------------------|-----------------|
|                                  |                   |                  |                        |                 | $\Delta G(B^-)^b$ | $\Delta G(BH)^c$ | $\delta\Delta G(BH)^d$ | $pK_{af}(BH)^e$ |
| 1c-1c1                           | -30.596           | -30.413          | -0.564                 | -0.413          | -30.586           | -30.299          | -0.418                 | -0.306          |
| 1c-1c2                           | -30.523           | -30.413          | -0.491                 | -0.360          | -30.472           | -30.299          | -0.304                 | -0.223          |
| 1c-1c3                           | -30.501           | -30.413          | -0.469                 | -0.344          | -30.494           | -30.299          | -0.326                 | -0.239          |
| 1c-1c4                           | -29.921           | -30.413          | 0.111                  | 0.081           | -29.920           | -30.299          | 0.248                  | 0.182           |
| 1c-1c5                           | -30.775           | -30.413          | -0.743                 | -0.545          | -30.819           | -30.299          | -0.651                 | -0.478          |
| 2ec-2ec1                         | -30.088           | -29.807          | -0.662                 | -0.485          | -30.105           | -29.799          | -0.437                 | -0.320          |
| 2ec-2ec2                         | -29.976           | -29.807          | -0.550                 | -0.403          | -29.946           | -29.799          | -0.278                 | -0.204          |
| 2ec-2ec3                         | -30.550           | -29.807          | -1.124                 | -0.824          | -30.504           | -29.799          | -0.836                 | -0.613          |
| 2ec-2ec4                         | -30.703           | -29.807          | -1.277                 | -0.936          | -30.344           | -29.799          | -0.676                 | -0.496          |
| 2ec-2ec5                         | -30.593           | -29.807          | -1.167                 | -0.856          | -30.614           | -29.799          | -0.946                 | -0.694          |
| 3cg-3cg1                         | -39.337           | -40.939          | 1.222                  | 0.896           | -39.148           | -40.939          | 1.661                  | 1.218           |
| 3cg-3cg2                         | -39.788           | -40.939          | 0.770                  | 0.565           | -39.475           | -40.939          | 1.334                  | 0.978           |
| 3cg-3cg3                         | -37.017           | -40.939          | 3.541                  | 2.597           | -36.997           | -40.939          | 3.812                  | 2.795           |
| 3cg-3cg4                         | -38.740           | -40.939          | 1.818                  | 1.333           | -38.754           | -40.939          | 2.055                  | 1.507           |

| 3cg-3cg5       | -38.085 | -40.939 | 2.473  | 1.814  | -38.028 | -40.939 | 2.780  | 2.039  |
|----------------|---------|---------|--------|--------|---------|---------|--------|--------|
| 3cg-3cg6       | -38.207 | -40.939 | 2.351  | 1.724  | -38.278 | -40.939 | 2.530  | 1.856  |
| 3cg-3cg7       | -38.212 | -40.939 | 2.347  | 1.721  | -38.243 | -40.939 | 2.566  | 1.881  |
| 4ecg-4ecg1     | -39.190 | -40.396 | 0.824  | 0.605  | -39.157 | -40.212 | 0.923  | 0.677  |
| 4ecg-4ecg2     | -38.540 | -40.396 | 1.475  | 1.081  | -38.551 | -40.212 | 1.530  | 1.122  |
| 4ecg-4ecg3     | -39.082 | -40.396 | 0.932  | 0.684  | -39.050 | -40.212 | 1.030  | 0.756  |
| 4ecg-4ecg4     | -39.371 | -40.396 | 0.643  | 0.472  | -39.333 | -40.212 | 0.748  | 0.548  |
| 4ecg-4ecg5     | -38.102 | -40.396 | 1.913  | 1.403  | -38.096 | -40.212 | 1.985  | 1.455  |
| 4ecg-4ecg6     | -39.592 | -40.396 | 0.423  | 0.310  | -39.443 | -40.212 | 0.638  | 0.468  |
| 4ecg-4ecg7     | -36.490 | -40.396 | 3.525  | 2.585  | -36.481 | -40.212 | 3.600  | 2.640  |
| 5egc-5egc1     | -31.579 | -31.986 | 0.027  | 0.020  | -31.557 | -31.985 | 0.297  | 0.218  |
| 5egc-5egc2     | -31.579 | -31.986 | 0.027  | 0.020  | -31.267 | -31.985 | 0.587  | 0.431  |
| 5egc-5egc3     | -30.489 | -31.986 | 1.117  | 0.819  | -30.505 | -31.985 | 1.349  | 0.990  |
| 5egc-5egc4     | -31.354 | -31.986 | 0.251  | 0.184  | -31.396 | -31.985 | 0.458  | 0.336  |
| 5egc-5egc5     | -31.548 | -31.986 | 0.057  | 0.042  | -31.552 | -31.985 | 0.302  | 0.221  |
| 5egc-5egc6     | -31.406 | -31.986 | 0.199  | 0.146  | -31.429 | -31.985 | 0.425  | 0.312  |
| 6egcg-6egcg1   | -41.002 | -42.786 | 1.403  | 1.029  | -41.177 | -42.902 | 1.594  | 1.169  |
| 6egcg-6egcg2   | -39.427 | -42.786 | 2.978  | 2.184  | -39.378 | -42.902 | 3.393  | 2.488  |
| 6egcg-6egcg3   | -39.062 | -42.786 | 3.342  | 2.451  | -39.020 | -42.902 | 3.751  | 2.751  |
| 6egcg-6egcg4   | -39.643 | -42.786 | 2.762  | 2.025  | -39.708 | -42.902 | 3.063  | 2.246  |
| 6egcg-6egcg5   | -39.540 | -42.786 | 2.864  | 2.100  | -39.549 | -42.902 | 3.222  | 2.363  |
| 6egcg-6egcg6   | -39.862 | -42.786 | 2.543  | 1.865  | -39.899 | -42.902 | 2.872  | 2.106  |
| 6egcg-6egcg7   | -40.516 | -42.786 | 1.889  | 1.385  | -40.181 | -42.902 | 2.590  | 1.899  |
| 6egcg-6egcg8   | -39.996 | -42.786 | 2.409  | 1.767  | -40.430 | -42.902 | 2.341  | 1.717  |
| 7ga-7ga1       | -23.642 | -24.380 | 0.357  | 0.262  | -23.630 | -24.367 | 0.606  | 0.444  |
| 7ga-7ga2       | -23.451 | -24.380 | 0.548  | 0.402  | -23.458 | -24.367 | 0.778  | 0.571  |
| 7ga-7ga3       | -24.236 | -24.380 | -0.237 | -0.174 | -24.207 | -24.367 | 0.030  | 0.022  |
| 7ga-7ga4       | -23.944 | -24.380 | 0.054  | 0.040  | -23.913 | -24.367 | 0.323  | 0.237  |
| 8caffa-8caffa1 | -25.001 | -25.071 | -0.312 | -0.229 | -24.989 | -25.069 | -0.051 | -0.037 |
| 8caffa-8caff2  | -24.448 | -25.071 | 0.242  | 0.177  | -24.441 | -25.069 | 0.498  | 0.365  |
| 8caffa-8caffa2 | -24.929 | -25.071 | -0.240 | -0.176 | -24.915 | -25.069 | 0.023  | 0.017  |

<sup>a</sup>: These symbols represent the protonation process for the molecules given in Fig.1

<sup>b</sup>  $\Delta G(B^-)$ : Formation Gibbs free energy of  $B^-$

<sup>c</sup>  $\Delta G(BH)$ : Formation Gibbs free energy of protonated  $BH$

<sup>d</sup>  $\delta\Delta G(BH)$ : Gibbs free energy calculated according to the Eq. 2 for the reaction (1)

<sup>e</sup>  $pK_a(BH)$ : Thermokinetic  $pK_a$  values calculated according to the Eq. 5

Table 5. pK<sub>a</sub> order belonging to the -OH group protons in the structure of molecules calculated from the different methods

| PM6 (directly calculated by MOPAC2009 using pK <sub>a</sub> keyword) |        |     |        |     |        |      |        |      |        |       |        |     |        |        |        |
|--|--------|-----|--------|-----|--------|------|--------|------|--------|-------|--------|-----|--------|--------|--------|
| 1c   |        | 2ec |        | 3cg |        | 4ecg |        | 5egc |        | 6egcg |        | 7ga |        | 8caffa |        |
| 1-H  | 8.644  | 1-H | 8.644  | 6-H | 7.881  | 6-H  | 7.881  | 2-H  | 8.415  | 6-H   | 7.832  | 2-H | 3.942  | 3-H    | 3.644  |
| 2-H  | 9.333  | 2-H | 9.333  | 2-H | 8.374  | 2-H  | 8.374  | 6-H  | 9.020  | 2-H   | 8.237  | 1-H | 8.865  | 2-H    | 9.276  |
| 5-H  | 10.748 | 5-H | 10.748 | 5-H | 9.170  | 5-H  | 9.170  | 1-H  | 10.584 | 4-H   | 8.857  | 3-H | 11.228 | 1-H    | 12.690 |
| 4-H  | 12.647 | 4-H | 12.647 | 1-H | 10.248 | 1-H  | 10.248 | 4-H  | 11.245 | 1-H   | 10.011 | 4-H | 14.817 |        |        |
| 3-H  | 14.089 | 3-H | 14.089 | 7-H | 11.417 | 7-H  | 11.417 | 3-H  | 13.942 | 8-H   | 10.590 |     |        |        |        |
|  |        |     |        | 4-H | 12.340 | 4-H  | 12.340 | 5-H  | 14.746 | 3-H   | 11.459 |     |        |        |        |
|  |        |     |        | 3-H | 13.138 | 3-H  | 13.138 |      |        | 7-H   | 13.011 |     |        |        |        |
|  |        |     |        |     |        |      |        |      |        | 5-H   | 13.735 |     |        |        |        |
| ΔG <sub>f</sub> (Thermokinetic)                                      |        |     |        |     |        |      |        |      |        |       |        |     |        |        |        |
| 1c   |        | 2ec |        | 3cg |        | 4ecg |        | 5egc |        | 6egcg |        | 7ga |        | 8caffa |        |
| 2-H  | 9.838  | 5-H | 12.199 | 7-H | 9.060  | 6-H  | 12.287 | 4-H  | 11.497 | 8-H   | 8.177  | 2-H | 8.512  | 3-H    | 8.134  |
| 1-H  | 9.930  | 4-H | 14.271 | 3-H | 12.255 | 3-H  | 12.616 | 1-H  | 12.442 | 7-H   | 12.134 | 1-H | 9.841  | 1-H    | 10.704 |
| 5-H  | 9.958  | 1-H | 12.539 | 2-H | 14.711 | 2-H  | 14.268 | 2-H  | 12.442 | 6-H   | 13.016 | 4-H | 9.957  | 2-H    | 14.809 |
| 4-H  | 12.863 | 2-H | 12.571 | 5-H | 14.777 | 4-H  | 14.429 | 6-H  | 13.190 | 5-H   | 14.244 | 3-H | 10.397 |        |        |
| 3-H  | 24.219 | 3-H | 26.263 | 6-H | 16.750 | 7-H  | 15.728 | 5-H  | 14.823 | 1-H   | 15.289 |     |        |        |        |
|  |        |     |        | 4-H | 18.426 | 5-H  | 16.607 | 3-H  | 27.579 | 2-H   | 15.951 |     |        |        |        |
|  |        |     |        | 1-H | 18.836 | 1-H  | 18.109 |      |        | 3-H   | 17.408 |     |        |        |        |
|  |        |     |        |     |        |      |        |      |        | 4-H   | 20.437 |     |        |        |        |
| ΔG (Thermodynamic)   |        |     |        |     |        |      |        |      |        |       |        |     |        |        |        |
| 1c   |        | 2ec |        | 3cg |        | 4ecg |        | 5egc |        | 6egcg |        | 7ga |        | 8caffa |        |
| 5-H  | -0.545 | 4-H | -0.936 | 2-H | 0.565  | 6-H  | 0.310  | 1-H  | 0.020  | 1-H   | 1.029  | 3-H | -0.174 | 1-H    | -0.229 |
| 1-H  | -0.413 | 5-H | -0.856 | 1-H | 0.896  | 4-H  | 0.472  | 2-H  | 0.020  | 7-H   | 1.385  | 4-H | 0.040  | 3-H    | -0.176 |
| 2-H  | -0.360 | 3-H | -0.824 | 4-H | 1.333  | 1-H  | 0.605  | 5-H  | 0.042  | 8-H   | 1.767  | 1-H | 0.262  | 2-H    | 0.177  |
| 3-H  | -0.344 | 1-H | -0.485 | 7-H | 1.721  | 3-H  | 0.684  | 6-H  | 0.146  | 6-H   | 1.865  | 2-H | 0.402  |        |        |
| 4-H  | 0.081  | 2-H | -0.403 | 6-H | 1.724  | 2-H  | 1.081  | 4-H  | 0.184  | 4-H   | 2.025  |     |        |        |        |
|  |        |     |        | 5-H | 1.814  | 5-H  | 1.403  | 3-H  | 0.819  | 5-H   | 2.100  |     |        |        |        |
|  |        |     |        | 3-H | 2.597  | 7-H  | 2.585  |      |        | 2-H   | 2.184  |     |        |        |        |
|  |        |     |        |     |        |      |        |      |        | 3-H   | 2.451  |     |        |        |        |
| Marvinbeans  |        |     |        |     |        |      |        |      |        |       |        |     |        |        |        |
| 1c   |        | 2ec |        | 3cg |        | 4ecg |        | 5egc |        | 6egcg |        | 7ga |        | 8caffa |        |
| 1-H  | 6.580  | 1-H | 5.974  | 1-H | 6.418  | 1-H  | 6.451  | 1-H  | 6.202  | 1-H   | 6.362  | 2-H | 1.674  | 3-H    | 2.345  |
| 5-H  | 7.523  | 4-H | 6.518  | 4-H | 6.707  | 6-H  | 6.574  | 4-H  | 7.164  | 6-H   | 6.948  | 3-H | 5.979  | 2-H    | 6.854  |
| 2-H  | 8.896  | 2-H | 7.531  | 7-H | 6.826  | 4-H  | 6.994  | 6-H  | 7.164  | 7-H   | 7.094  | 4-H | 7.764  | 1-H    | 7.986  |
| 4-H  | 9.278  | 5-H | 8.790  | 6-H | 6.991  | 3-H  | 7.011  | 5-H  | 8.070  | 8-H   | 7.161  | 1-H | 7.866  |        |        |
| 3-H  | 12.364 | 3-H | 12.404 | 3-H | 7.139  | 7-H  | 7.277  | 2-H  | 8.134  | 4-H   | 7.173  |     |        |        |        |
|  |        |     |        | 5-H | 8.842  | 5-H  | 8.773  | 3-H  | 12.312 | 3-H   | 7.183  |     |        |        |        |
|  |        |     |        | 2-H | 8.897  | 2-H  | 9.035  |      |        | 5-H   | 8.124  |     |        |        |        |
|  |        |     |        |     |        |      |        |      |        | 2-H   | 9.013  |     |        |        |        |
| Sparc  |        |     |        |     |        |      |        |      |        |       |        |     |        |        |        |
| 1c   |        | 2ec |        | 3cg |        | 4ecg |        | 5egc |        | 6egcg |        | 7ga |        | 8caffa |        |
| 4-H  | 8.790  | 4-H | 8.790  | 6-H | 6.690  | 6-H  | 6.690  | 4-H  | 8.610  | 6-H   | 6.690  | 2-H | 3.980  | 3-H    | 4.400  |

Table 5. (Continued) pK<sub>a</sub> order belonging to the -OH group protons in the structure of molecules calculated from the different methods

|     |        |     |        |     |       |     |       |     |        |     |       |     |       |     |       |
|-----|--------|-----|--------|-----|-------|-----|-------|-----|--------|-----|-------|-----|-------|-----|-------|
| 2-H | 9.090  | 2-H | 9.090  | 3-H | 7.980 | 3-H | 7.980 | 6-H | 8.610  | 3-H | 7.980 | 4-H | 6.910 | 1-H | 7.890 |
| 5-H | 9.360  | 5-H | 9.360  | 7-H | 7.980 | 7-H | 7.980 | 5-H | 8.840  | 7-H | 7.980 | 1-H | 8.590 | 2-H | 9.070 |
| 1-H | 9.680  | 1-H | 9.680  | 2-H | 8.550 | 2-H | 8.550 | 2-H | 8.990  | 4-H | 8.380 | 3-H | 8.590 |     |       |
| 3-H | 14.900 | 3-H | 15.040 | 5-H | 8.560 | 5-H | 8.560 | 1-H | 9.680  | 8-H | 8.380 |     |       |     |       |
|     |        |     |        | 4-H | 9.280 | 4-H | 9.280 | 3-H | 15.220 | 2-H | 8.440 |     |       |     |       |
|     |        |     |        | 1-H | 9.500 | 1-H | 9.500 |     |        | 5-H | 8.760 |     |       |     |       |
|     |        |     |        |     |       |     |       |     |        | 1-H | 9.500 |     |       |     |       |

### 3. COMPUTATIONAL METHOD

The CS Chemoffice and the MOPAC2009 programs were used for the calculation of the physicochemical properties of the tea components. All the molecules were drawn by the CS ChemDraw and then minimized by the MM2 method in the CS Chem 3D program (MOPAC 2009). Their theoretical calculations in aqueous and blood phases were carried out using the new semi-empirical PM6 quantum chemical methods in the MOPAC2009 program. The pK<sub>a</sub> values of the molecules were computed by MarvinSketch v5.3.1 (ChemAxon) and SPARC v4,5 (SPARC 2009) programs in aqueous and blood phases and the results were compared with those of experimental results.

The most stable conformations of all of the studied molecules were computed by all the present computational methods.

### 4. CONCLUSION

It is thought that not only the number of -OH group but also the geometry of the tea components has an important role in the reactivities of these compounds in human metabolism. Dipole moment and nucleophilicity did not give any important information from the viewpoint of the reactivity for these molecules. The best correlations could be observed between PM6 and experimental pK<sub>a</sub> values for **6egcg** and **8caffa**; PM6-ΔG<sub>f</sub> for **4ecg**; Marvin-experimental for **1c** and **2ec**; ΔG<sub>f</sub>-Sparc for **3cg**, ΔG<sub>f</sub>-ΔG for **5egc** and Marvinbeans-Sparc for **7ga** in the aqueous phase; and Marvinbeans-experimental for **1c** and **2ec**; ΔG<sub>f</sub>-Marvinbeans for **3cg**; ΔG-experimental for **4ecg**; ΔG<sub>f</sub>-Sparc for **5egc**; ΔG-Sparc for **6egcg**; and Marvinbeans-Sparc for **7ga** and ΔG<sub>f</sub>-experimental for **8caffa** in the blood phase.

Positive meaningful correlations ( $R^2 > 0.9999$ ) were observed between thermokinetic pK<sub>a</sub> values in both aqueous and blood phases for **1c**, **2ec**, **3cg**, **4ecg**, **5egc**, **6egcg** and **7ga**. It can be resulted that the mentioned molecules have showed similar behaviors like permeability in the blood phase with those molecules in the aqueous phase due to their good correlations.

A good correlation could not be observed among the pK<sub>a</sub> values for the similar acidic proton belonging to the -OH in the molecules for all calculation methods [PM6, ΔG<sub>f</sub> (Thermokinetic), ΔG (Thermodynamic), Marvinbeans and Sparc].

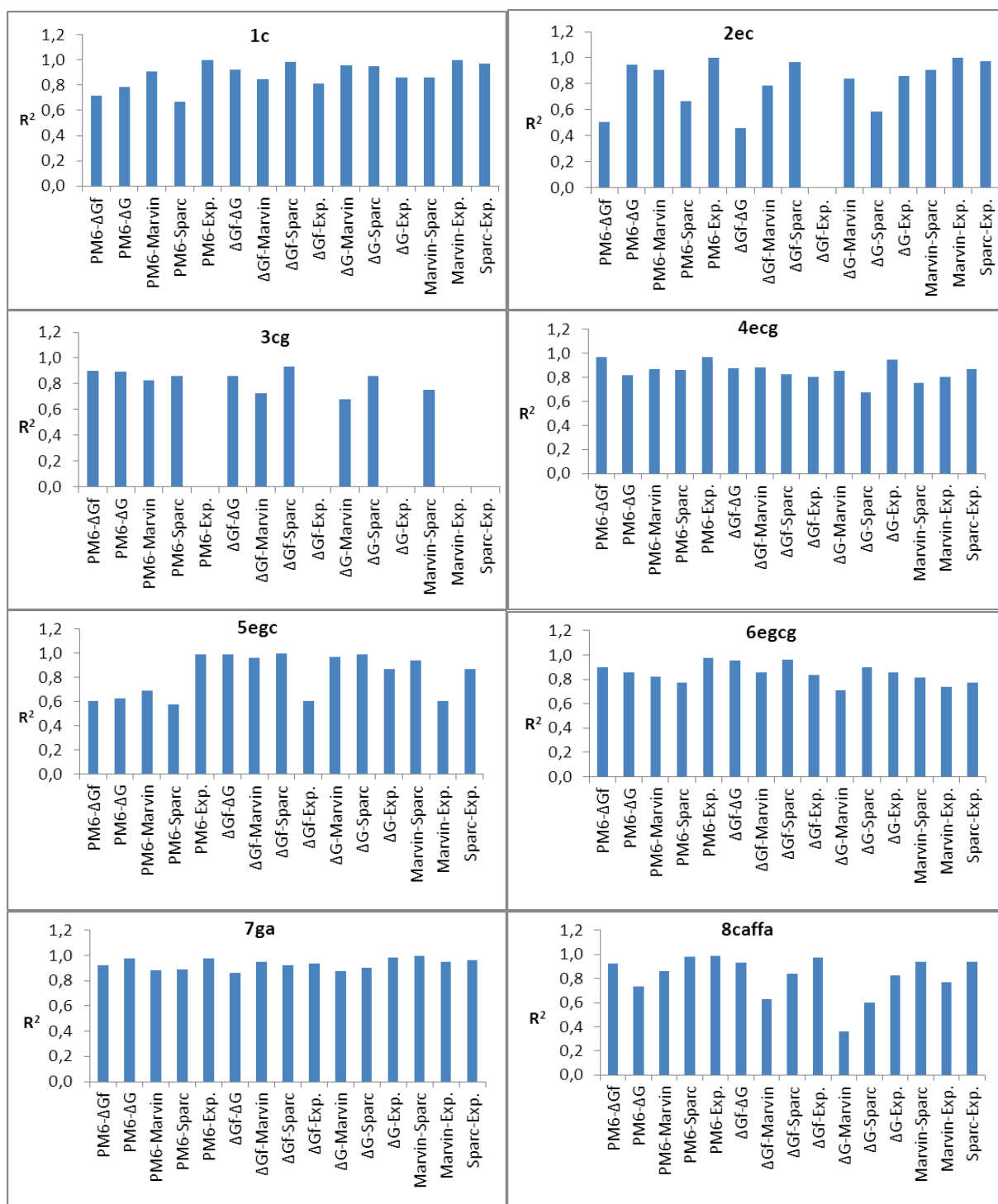


Figure 2. The correlations for  $pK_a$  values in the aqueous phase by different calculation methods [experimental: Tama et al. 2001; Adams et al. 2002; Inoue et al. 2002; Herrero-Martínez et al. 2005), PM6,  $\Delta G_f$  (Thermokinetic),  $\Delta G$  (Thermodynamic), Marvinbeans and Sparc],  $R^2$ : Correlation coefficient.

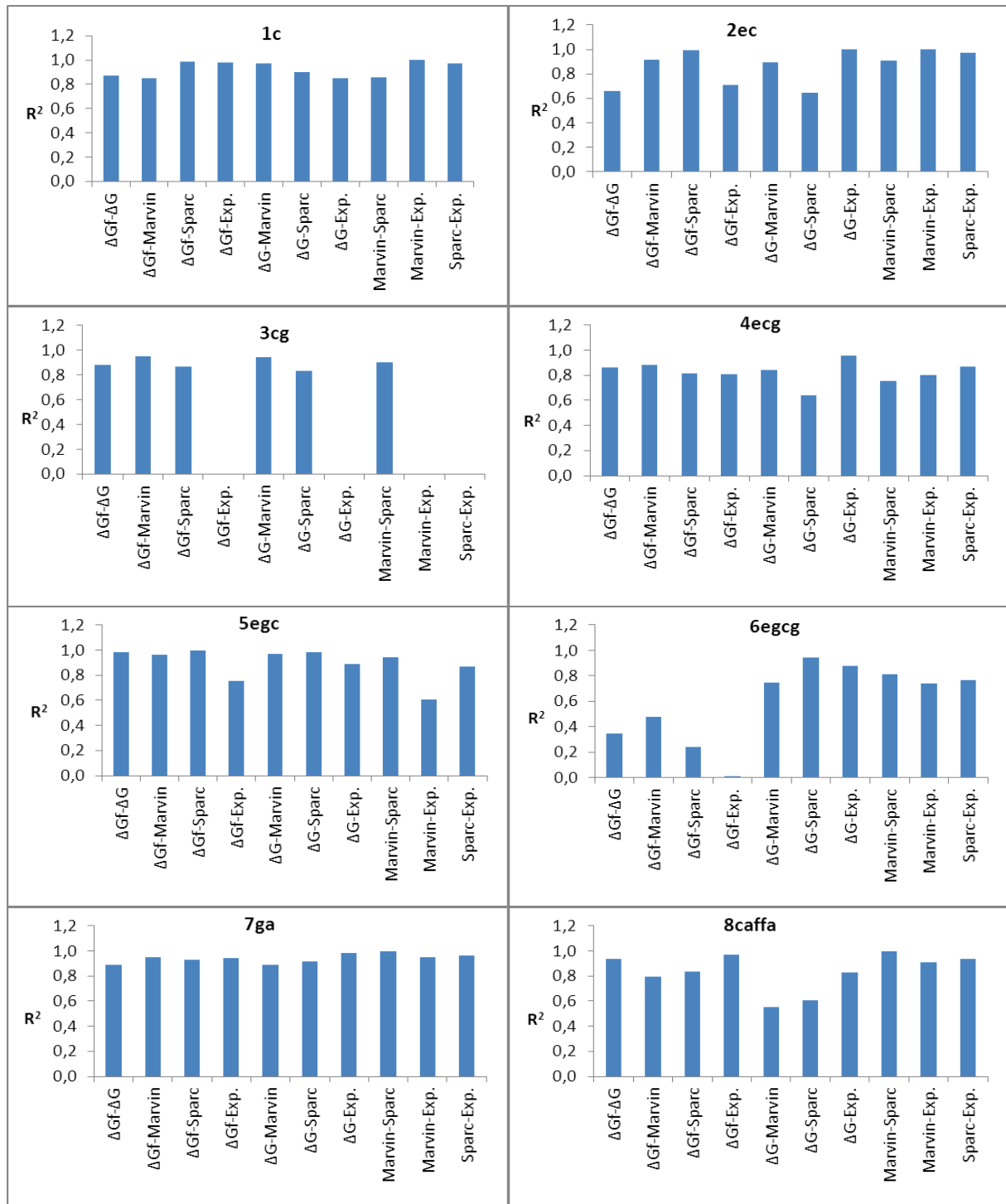


Figure 3. The correlations for  $pK_a$  values in the blood phase by different calculation methods [experimental: Tama et al. 2001; Adams et al. 2002; Inoue et al. 2002; Herrero-Martínez et al. 2005), PM6,  $\Delta G_f$  (Thermokinetic),  $\Delta G$  (Thermodynamic), Marvinbeans and Sparc],  $R^2$ : Correlation coefficient.



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